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Investigations on the drug releasing mechanism from an asymmetric membrane-coated capsule with an in situ formed delivery orifice

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Abstract

Asymmetric membrane-coated capsules with in situ formation of a delivery orifice were examined for their improved osmotic effects. The release mechanisms were investigated for drugs with both moderate to high water solubility and those with poor water solubility. The capsule wall membrane was produced by a phase-inversion process, in which an asymmetric membrane was formed on stainless steel mold pins by dipping the mold pins into a coating solution containing a polymeric material followed by dipping into a quenching solution. In situ formation of a delivery orifice in the thin membrane was proven by visualization of a jet stream of chlorophyll being released from the capsule. The release mechanism for drugs with moderate to high water solubility was mainly controlled by the osmotic effect, which is a function of the drug's solubility.
Permeability across the asymmetric membrane of the capsule was determined to be 4.28×10^{-6} c with water solubilities in a moderate to high range. Accordingly, the poorly water-soluble drug, nifedipine, was unable to create enough of an osmotic effect to activate drug release. Solubilization either by the addition of the solubility enhancer, SLS, or by a solid dispersion with HPMC could increase the solubility of nifedipine to a sufficient extent to activate drug release. It was found that the suspending ability induced by the viscous nature of HPMC further interacted with SLS to synergistically increase the maximal percent release and the release rate of nifedipine. The osmotic effect of this suspension ability was proposed as the underlying mechanism responsible for the release of poorly water-soluble drugs, i.e. nifedipine, from this system.

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Keywords: Asymmetric membrane; Osmotic pressure; Cellulose acetate; Nifedipine; In situ formation

1. Introduction interest in the development of osmotic devices over the past two decades. Designs of various types of It is known that pharmaceutical agents can be osmotic pumps have been reported $[1-3]$ and redelivered in a controlled pattern over a long period viewed [4,5]. Osmotic tablets with an asymmetric by osmotic pressure. There has been increasing membrane coating, which can achieve high water fluxes, have been described [6]. The asymmetric ***Corresponding author. Tel./fax: ¹886-2-2377-1942. membrane capsule described [7,8] is also an example *E-mail address:* hsiuoho@tmu.edu.tw (H.-O Ho). of a single-core osmotic delivery system consisting

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asymmetric membrane. One of the advantages of the some for commercial production. Although it is asymmetric membrane capsule is the higher rate of known that delivery orifices can be created in situ in water influx, allowing the release of drugs with semipermeable membranes, it is still necessary to lower osmotic pressures or lower solubilities. In spite utilize expanding material in the core of or added of this advantage, there are many instances where the pore formers in the membrane. Since asymmetric solubility of a drug is too low to provide a reason- membranes in osmotic membranes consist of a very able driving force for water ingress. In such situa- thin, dense skin structure supported by a thicker, tions, various methods have been reported to en- porous structural layer, in situ formation of delivery hance this driving force either by improving drug orifices on this thin layer is potentially possible with solubility, including the use of crystal habit modifiers no assistance. In this study, we explored whether [9], assisting with lyotropic crystals [10,11], adding delivery orifices could be formed in situ on pH-regulating excipients [12,13], complexing with asymmetrically coated membranes. Compliance with inclusion compounds [14], or by enhancing the an osmotic control mechanism of drug release and contact surface area of the drug by utilizing wicking the influential factors were investigated. agents [15].

In a majority of cases, osmotic delivery systems contain at least one delivery orifice in the membrane **2. Experimental section** for drug release. Pre-formation and in situ formation are two possible ways of creating delivery orifices in 2 .1. *Materials* membranes. Laser drilling is one of the most commonly used techniques to create preformed delivery Cellulose acetate (CA 398-10) was supplied by orifices in osmotic tablets [16]. The use of modified Eastman Chemicals Co. (Kingsport, USA). punches for producing a preformed delivery orifice Nifedipine (NF) and felodipine (FL) were provided in osmotic dosage forms is also possible [17]. by Merck (Darmstadt, Germany) and the Sigma Controlled-porosity osmotic pumps (CPOP), contain Chemical Co. (St Louis, MO, USA), respectively. As water-soluble additives in the coating membrane, nifedipine was light sensitive, all samples were kept which after coming in contact with water, dissolve in an amber-colored container, wrapped in aluminum resulting in in situ formation of a microporous foil, or covered by a blanket during the whole membrane [18–20]. The resulting membrane is experimental process. Chlorpheniramine maleate, substantially permeable to both water and dissolved pyridoxine, theophylline, chlorophyllin, glycerin, solutes. Because of that, the mechanism of drug Tween 80, sodium lauryl sulfate (SLS) were from release from these systems was found to be primarily the Sigma Chemical Co. (St Louis, MO, USA). osmotic, with simple diffusion to a minor extent. Triethyl citrate (TEC), acetonitrile, methanol were Systems with passageways formed in situ are also from Merck (Germany). Hydroxypropylmethylcelludescribed in US patent no. 5,736,159 [21]. A small lose (HPMC, 5 cps, 15 cps and 50 cps) was opening is formed at the edge of the tablet caused by purchased from the Shin-Etsu Chemical Co. (Japan). the expansion of expandable material in the core.

The size of the delivery orifice must be optimized 2.2. Methods in order to control the amount and rate of drug released from osmotic systems. Preformed delivery 2 .2.1. *Capsule preparation* orifices would be advantageous for the design of Capsules with asymmetric membranes were prodelivery orifices and hence the ability to control drug duced using a dip-coating process. The stainless steel release. However, a complicated continuous process mold pins were dipped into polymer solutions consuch as laser drilling, or specially designed machin-
sisting of 15% w/v cellulose acetate (CA 398-10, ery such as modified punches would be required for Eastman Fine Chemical) dissolved in a mixture of commercial production scale. Therefore, in situ acetone/alcohol/glycerin (62 ml/34.5 ml/10 g), formation of delivery orifices in the semipermeable followed by quenching in an aqueous solution (10%

of a drug-containing core surrounded by an membrane of osmotic systems would be less burden-

w/v glycerin). After quenching, the pins were with- 2 .2.5. *HPLC analysis* drawn and allowed to air-dry. Then, the capsules The HPLC system consisted of a Rainin solvent were stripped off the pins, trimmed to size, and kept delivery pump (Dynamax, model SD-200), a UV in desiccators until use. detector (Dynamax, model UV-1), an automatic

nifedipine and sodium lauryl sulfate with hand shaking in a plastic bag for at least 15 min. A solvent method was employed to prepare solid dispersion 2.3. *Theoretical considerations* systems for nifedipine. HPMC were selected as the water-soluble polymers. After dissolving nifedipine and HPMC in a suitable volume of an acetone/water of an acetone in a suitable volume of an acetone water mixture, the so ture of 50–60 °C. Dried residues were ground with a rated in a forced-air convention oven at a tempera-
ture of 50–60 °C. Dried residues were ground with a
coffee mill, and granules passing an 80-mesh sieve where dV/dt is the volumetric influx rate of water
then stored in desiccators protected from light until
use. After the filling operation, the capsules were
capped and sealed with a sealing solution, which
contains 16% ce

2.2.3. Solubility tests

Excess drugs were suspended in deionized water and maintained at 37° C for at least 72 h with

An in vitro dissolution test was performed using USP dissolution methodology (Apparatus 2, 50 rpm, 37 °C, 500–1000 ml of medium with sinker) (JASCO, Model DT-610). In all cases, an appro- where *R* is the universal gas constant, *T* is the priate volume of sample was withdrawn at pre- temperature, *M*.*W*. is molecular weight, and *S* is the determined time intervals and assayed by either a saturation solubility of the single component (drug). validated UV absorbance measurement or by an Substituting for $\Delta \pi$ into Eq. (1) and substituting the HPLC/UV method (Helios, Unicam and Dynamax, resultant expression for d*V*/d*t* into Eq. (2), the Rainin Instrument). The same state of the state of th

sample injector (Dynamax, model AI-3), and an 2.2.2. Osmotic pump capsule preparation

Asymmetric membrane capsules were fabricated

and filled with the desired amount of drug or drug-

excipient mixture by hand. Physical mixtures of

in a ratio of 3:7 (v/v). A flow-

$$
\frac{dV}{dt} = \frac{A}{h} L_p \sigma \Delta \pi \tag{1}
$$

the release profile is given by:

$$
\frac{dM}{dt} = \frac{dV}{dt}S\tag{2}
$$

intermittent shaking. Immediately after filtration where dM/dt is the release rate, dV/dt is given by
from the syringe, filtrate in the middle portion was
sampled and properly diluted. The drug concen-
tration was assay on one side of the capsule wall and sink conditions (assumed) outside the capsule walls. Also, assuming 2.2.4. *Release test* ideality, the expression for $\Delta \pi$ can be written as:

$$
\Delta \pi = MRT = \frac{S}{M.W.}RT\tag{3}
$$

$$
\frac{dM}{dt} = \left(\frac{A}{h}L_p \sigma RT\right) \frac{S^2}{M.W.}
$$
\n(4)

Eq. (4) indicates that a plot of the release rate versus (S^2 /*M*.*W*.) should be linear with a slope given by the expression in parentheses. Based on Eq. (4), the water permeability (L_p) of the asymmetric membrane capsule wall was calculated.

3. Results and discussion

The asymmetric membrane-coated capsules prepared appeared to be white, opaque, and glossy with no visible imperfections. Weight variations in the asymmetric membrane capsules and their dimensions were demonstrated to be consistent with little variation. This confirms that the process of producing these capsules is reproducible. Scanning electron micrographs (SEMs) of the capsule walls show that the membrane was asymmetric with a relatively thin dense region on a porous substrate with longer micropores (Fig. 1). No pore structures were shown in the dense region (Fig. 1A). The porous region at both \times 100 (Fig. 1B) and \times 200 magnification (Fig. 1C) reveals numerous pore structures.

In situ formation of a delivery orifice for releasing drug was proven with photographs as shown in Fig. 2 in which a deeply colored jet stream of chlorophyll from an open hole can be observed when an asymmetric membrane-coated capsule encapsulated with chlorophyll was suspended in the water medium. This delivery process continued for another 30 min as demonstrated by Fig. 2B. However, when this capsule was suspended in a 5% NaCl solution, the osmotic effect was inactivated, and no release of chlorophyll (Fig. 2C) was observed. This indicates that in situ formation of a delivery orifice is possible in the thin structure of the asymmetric membrane. The osmotic pressure created might play an important role in the switching on or off of this mechanism.

The osmotically controlled drug release mechanism from asymmetric membrane-coated capsules Fig. 1. Scanning electron micrographs of asymmetric membrane was further characterized based on Eq. (4) using capsule wall at formulation A (A) dense region (outer layer) at drugs of varying solubilities. The core formulation $\times 2000$ magnification, (B) cross-section at $\times 100$ magnification, consisted of drug alone but with varying solubilities (C) porous region (inner layer) at \times 200 magnification.

 (A)

 (B)

Fig. 2. Photograph of asymmetric membrane capsule with chlorophyllin in water. (A) After 30 min, (B) After 90 min, (C) In NaCl 5% solution.

theophylline, felodipine, and nifedipine); these were asymmetric membrane capsules are shown in Fig. 3. individually loaded into the asymmetric membrane The initial portion of the drugs release profiles were

in water (chlorpheniramine maleate, pyridoxine, capsule. The in vitro drug release profiles from

Fig. 3. Release profiles of drugs from asymmetric membrane capsule (formulation A) in water (50 rpm, $n=3$). Key: (\bullet) Chlorpheniramine maleate 50 mg; (\heartsuit) pyridoxine 50 mg; (∇) theophylline 50 mg; (∇) felodipine 50 mg; (\blacksquare) nifedipine 50 mg.

Drug	M.W.	$S \text{ (mg/ml)}$	$S^2/M.W.$	Release rate $(\frac{9}{6}$ /h)	Release rate (mg/h)
Chlorpheniramine maleate	390	576.72 ± 16.24	852.84	33.90	16.95
Pyridoxine	205	224.47 ± 0.96	245.79	12.58	6.29
Theophylline	180	7.18 ± 0.03	0.29	1.30	0.65
Felodipine	384	\leq 1	~ 0	\sim 0	\sim 0
Nifedipine	346	\leq 1	~ 0	\sim 0	\sim 0

Table 1 The solubility of drugs in water at 37 °C ($n=3$)

used to calculate the initial drug release rate. The initial portion of the profiles increased linearly with results in Fig. 3 reveal that both the amount released respect to the square of the drug solubility divided by and the release rate were a function of drug solu- the molecular weight of the corresponding drug as bility. In comparison with the drug solubilities listed predicted by Eq. (4). A statistically significant in Table 1, the amount released was larger and the correlation of $r^2 = 0.9936$ was demonstrated. This drug release rate was faster as drug solubility complies with the drug released by an osmotic increased. pumping mechanism. Assuming ideality, where *R* is

release rate (calculated from the slope of the drug comparable to that from a similar membrane design
release profile) and $S^2/M.W$. was observed. The slope reported in the literature [8].
of linear portion is 0.0185 cm h all other factors constant, the drug release rate in the capsules with an in situ formed delivery system, the

The graph displayed in Fig. 4 plots the release rate the Universal Gas Constant and T is the absolute (dM/dt) versus the ratio of the square of drug temperature, L_p (at 37 °C) is calculated to be 4.28 ×
solubility to molecular weight $(S^2/M.W.)$ based on 10^{-6} cm²/h-atm based on Eq. (4) with known values
Eq. (4). A lin

Fig. 4. Linear relationship between the release rate and the square of solubility divided by molecular weight of the drug.

osmotic effect is the main activation force for drug Table 2
release Similarly drug solubility is expected to be The solubility of nifedipine in SLS solution at 37 °C (n=3) release. Similarly, drug solubility is expected to be the determining factor for the success of engineering asymmetric membrane-coated capsules with an in situ formed delivery orifice with a desirable release rate. It is likely that a drug with low solubility would not create enough osmotic pressure to activate drug release. Because of this, the drug release mechanism from an asymmetric membrane-coated capsule with in situ formation of a delivery orifice was further studied by examining the influence of core formulation variables including the added amount and viscosity of hydroxypropylmethylcellulose (HPMC) apparently increased with the increased amount of and the amount of sodium lauryl sulfate (SLS). added SLS. Possibly, the greater the amount of SLS Nifedipine was selected as a model drug because it is which is incorporated into the capsule, the larger has poor water solubility. All subsequent release the osmotic effects will be which can be activated to studies were done in water medium with the addition cause a greater amount of nifedipine to be dissolved of 1% Tween 80 as the solubilizing agent. and released. The osmotic effects of SLS on the

formulation had a marked influence on nifedipine could be attributed to two factors. One is the release. When the added amount of SLS was at a 1:1 solubilization effect of SLS (Table 2) in enhancing ratio to the nifedipine amount, only 2% of the the dissolved amount of nifedipine in the core nifedipine was released, whereas the release amount medium for increasing the osmotic effect, and the of nifedipine increased to 40% by adding SLS at a other is that the dissolved SLS acts as an osmotic 20:1 ratio to the nifedipine amount. The release rate agent to increase the osmotic effect. Since SLS was

SLS concentration $(w/v\%)$	Solubility (mg/ml)	S.D.
0.1	0.021	0.0002
0.5	0.166	0.0028
1.0	0.371	0.0063
2.5	0.799	0.0092
5.0	1.362	0.0099
10.0	2.296	0.0104
20.0	3.280	0.0425

Fig. 5 shows that the amount of SLS in the core release rate and the released amount of nifedipine

Fig. 5. Release profiles of nifedipine from asymmetric membrane capsule in 1% Tween 80 solution (50 rpm, $n=3$). Nifedipine (NF) was made by physically mixed method with SLS (S). NF/S ratio: (\bullet) 1/1, (\circ) 1/5, (∇) 1/10, \circ) 1/20.

was released in company with nifedipine. When the with SLS. Compared with the core formulation SLS was exhausted, both mechanisms ceased, which which only contains nifedipine and SLS at a ratio of terminated the release of nifedipine from these 1:10, the addition of HPMC of varying viscosities capsules. This leads to the released amount of further increased both the release rate and the nifedipine being proportional to the added amount of released amount of nifedipine. At the same level of SLS. This quantitative relationship is illustrated in HPMC, an increase in the released percentage of Fig. 6 by plotting the maximal percent released 60% was shown for HPMC with a viscosity of 5 cps, versus the ratio of SLS to nifedipine. The slope whereas increases to 70–80% were observed for (2.196) of the linear plot can be used to predict what HPMC with a viscosity of either 15 or 50 cps. The added amount of SLS per unit amount of nifedipine further promotion of the released amount of would be necessary to reach a maximal 100% release infedipine by all viscosity grades of HPMC could be of nifedipine from this capsule system. Based on attributed to the enhancement of nifedipine solubility this, quite a large amount (about 440 mg) of SLS with the aid of the solid dispersion. However, this was possibly needed to completely release 10 mg seems to indicate that the higher the viscosity of nifedipine from this type of capsule by extrapolation. HPMC used, the larger amount of nifedipine which An in vitro dissolution test of nifedipine mixed with could be released and the faster the release rate sodium lauryl sulfate at a weight ratio of 1:44 had which would result. Since the enhancement of been performed. It was fairly confirmed that the nifedipine solubility using various grades of HPMC extent of drug release from this formulation was is comparable (Table 3), another mechanism seems increased to about 85%. The result shown is closely to be operating to have such an influence. consistent with the extrapolated prediction. Fig. 8 demonstrates the effect of different added

HPMC at the same level on the release pattern of cps) on the release pattern of nifedipine. The added nifedipine from these osmotic capsules with an in amount of HPMC also had a pronounced influence situ formed delivery orifice. Nifedipine was incorpo- on the release profile. Since an improvement in rated with HPMC in a solid dispersion form prepared solubility of nifedipine using different ratios of

released through the in situ formed delivery orifice, it by the solvent method and then physically mixed

Fig. 7 illustrates the effect of viscosity grades of amounts of HPMC of the same viscosity grade (50

Fig. 6. Correlation of max released and SLS/nifedipine ratio.

Fig. 7. Release profiles of nifedipine in 1% Tween 80 solution (50 rpm, $n=3$). Nifedipine (NF) was made by solvent method with HPMC and physically mixed with SLS (S). Key: (\bullet) NF/S ratio: $1/10$; (\circ) NF/HPMC 5 cps/S ratio: $1/10/10$; (\bullet) NF/HPMC 15 cps/S ratio: $1/10/10$; (∇) NF/HPMC 50 cps/S ratio: $1/10/10$.

HPMC plays as a thickening agent in elevating the of HPMC of varying viscosity grades. That is, a viscosity of the core suspension and, subsequently, higher-viscosity HPMC would promote the morepreventing precipitation of nifedipine particles was efficient suspension of nifedipine particles for dissosuspected of possibly being the expression of a larger lution, leading to an increase in the released amount surface for dissolution. The larger the amount of with an increase in the viscosity grade of HPMC HPMC used, the higher the viscosity of the core used for preparing the solid dispersion. suspension would be, leading to the efficient suspen- Based on the above release profiles, we concluded sion of nifedipine particles in the capsule. As a that this asymmetric membrane-coated capsule with consequence, the release rate increased when in- an in situ formed delivery orifice was able to release creasing the added amount of HPMC by increasing a water-insoluble drug such as nifedipine in the the available surface area for dissolution. This mech- presence of an osmotic agent with the aid of a

The solubility of various forms of nifedipine in water at 37 °C

Formulation	Ratio	Solubility (mg/ml)
Nifedipine		0.011^{a}
Nifedipine/HPMC 5 cps	1:10	0.0422 ± 0.0010
Nifedipine/HPMC 15 cps	1:10	0.0463 ± 0.0021
Nifedipine/HPMC 50 cps	1:10	0.0439 ± 0.0021
Nifedipine/mannitol	1:5	0.0132 ± 0.0001
Nifedipine/mannitol	1:10	0.0104 ± 0.0003

 a From Ref. [22].

HPMC to nifedipine was not obvious, the role anism seems capable of being explained by the effect

solubilizing agent and a suspension agent. Therefore, Table 3
the system was operated by an osmotic-suspension
The solubility of various forms of pifedining in water at 37 °C co-controlled delivery mechanism somewhat differ- $(n=3)$ ent from either the generic EOP or the push-pull osmotic tablet. This proposed mechanism was further supported by the results demonstrated in Fig. 9.

> Fig. 9A shows that the release of nifedipine was activated with a 1-h delay from the time the capsules were filled with the physical mixture of nifedipine and SLS at a ratio of 1:10. Two hours later, equilibrium had been reached with a release rate of about 0.5 mg/h, after which the release rate declined

Fig. 8. Release profiles of nifedipine in 1% Tween 80 solution (50 rpm, $n=3$). Nifedipine (NF) was made by the solvent method with HPMC 50 cps (H) and physically mixed with SLS (S). NF/H/S ratio: (\bullet) 1/1/10, (\circ) 1/5/10, (∇) 1/10/10.

almost completely ceased by 10 h. In the presence of of nifedipine continued for more than 24 h with an osmotic agent such as SLS, water imbibed by the nifedipine in a solid dispersion form with HPMC. semi-permeable membrane into the capsule was The release rate gradually reached its maximum at 8 gradually saturated with SLS to further build up the h and was maintained at a plateau until 20 h. osmotic pressure difference between the internal Compared to SLS, a longer period of release time system and the external environment. Simultaneous- but a lower release rate of nifedipine with HPMC ly, the dissolved SLS caused the solubilization of was demonstrated. A longer period of release nifedipine which was then available for release. occurred because it takes time for all the HPMC to During this period, the in situ formed delivery orifice completely dissolve in such quiescent conditions might be created at the weakest point in the inside the capsule, and the longer-sustained osmotic asymmetric membrane with an increase in osmotic effect of HPMC might be a result of it being too pressure. It is expected that there is a lag time to large in size to be released. A lower release rate of reach such a state, which means that the release of nifedipine with HPMC might be attributed to a lower drug is only activated with a 1-h lag time as shown solubilization effect of HPMC which resulted in a in Fig. 9A for SLS. After that time, dissolved reduced osmotic effect. However, maintaining a nifedipine and SLS were delivered through the maximal release rate of 0.25 mg/h with a drug orifice which had formed in situ. However, with the solubility of 43.9 μ g/ml (refer to Table 3) requires release of nifedipine at the expense of SLS, the an osmotic pressure higher than 7.8×10^3 atm acosmotic effect gradually diminished and the solubili- cording to Eq. (4). The induction of osmotic pressure zation effect was also retarded. This led to the by the presence of HPMC was determined to be release rate of nifedipine gradually decreasing with minimal (data not shown), since osmotic pressure is time. Then the release of nifedipine came to a a colligative property that is determined by the complete stop when all the added amount of SLS molecular number, which would be less for such a

4 h afterward, and the release of nifedipine had Nevertheless, Fig. 9A also shows that the release was exhausted. **polymer** as HPMC with a high molecular weight.

Fig. 9. Release rate of nifedipine from asymmetric membrane capsule in 1% Tween 80 solution (50 rpm, $n=3$). (A) Nifedipine (NF) was made by physically mixed with SLS (S) or solvent method with HPMC 50 cps (H). Key: (\bullet) NF/S ratio: 1/10; (O) NF/H ratio: 1/10. (B) Nifedipine (NF) was made by the solvent method with mannitol (M) and physically mixed with SLS (S). NF/M/S ratio: (\bullet) 1/10/10, (\circ) 1/5/10.

Therefore, the exact mechanism causing such a nifedipine being released in both its insoluble and release rate with such a drug solubility was not due soluble forms, the latter of which stands for the exact to the induction of a greater osmotic pressure. solubility. Because of that, with the same osmotic effect as in Fig. 9B further shows the release pattern of the case of SLS only, there must necessarily be a nifedipine from an encapsulated mixture consisting larger excess value in the solubility term than the of a solid dispersion of nifedipine with mannitol real solubility to have such a large release rate and a prepared by the solvent method and physically mixed cumulative released amount. This large excess value with SLS. A maximal amount of 20% was released in the solubility term could be attributed to for both mixtures, but with a faster rate for a larger

amount of mannitol. Mannitol is known for being an **4. Conclusions** osmotic agent which should induce osmotic pressures proportional to its amounts which will The in situ formed delivery orifice in controlledsynergize with the osmotic effect of SLS. This release polymeric capsules with an asymmetric memreveals that the higher osmotic effects induced by a brane wall is mainly responsible for the delivery of larger amount of mannitol with the same amount of both soluble and poorly soluble drugs. In vitro SLS does lead to an increase in the release rate, and release studies indicate that drug delivery from this it is maintained as it is until both are exhausted. asymmetric membrane-coated system is principally Nevertheless, it is the solubility that determines the controlled by osmotic pressure for those drugs with cumulative released amount of nifedinine during the moderate to high water solubilities. The asymmetric cumulative released amount of nifedipine during the moderate to high water solubilities. The asymmetric active period of osmotic pressure as predicted by E_0 membrane-coated capsule prepared in this study has active period of osmotic pressure as predicted by Eq. membrane-coated capsule prepared in this study has (4). Since the increase in nifedipine solubility with a permeability of 4.28×10^{-6} cm²/h-atm. This patro-fold two-fold increase in the amount of mannitol was rameter can be used to predict the release rate of any two-fold increase in the amount of mannitol was drug encapsulated in this asymmetric membrane insignificant as shown in Table 1, similarity in the drug encapsulated in this asymmetric membrane
maximal amount released is expected for both mix-
capsule. Solubilization of poorly water-soluble drugs maximal amount released is expected for both mix-
tures under a reasonable assumption that the active with the incorporation of solubility enhancers is able

pended. During the experiment, water was imbibed creating a viscous suspension in situ inside the **References** capsule, which resulted from the thickening agent (HPMC) and the imbibed water. Both the insoluble [1] F. Theeuwes, Elementary osmotic pump, J. Pharm. Sci. 64 and soluble forms of nifedipine in the suspension (1975) 1987–1991. were subsequently pumped out through the in situ [2] D.R. Swanson, B.L. Barclay, P.S.L. Wong, F. Theeuwes, formed delivery orifice. This explains why only the Nifedipine gastrointestinal therapeutic system, Am. J. Med. soluble form of nifedipine was released in formula-

⁸³ (Suppl. 6B) (1987) 3–9.

⁸³ (Suppl. 6B) (1987) 3–9.

⁸³ (Suppl. 6B) (1987) 3–9.

⁸³ (Suppl. 6B) (1987) 3–9. tions containing mannitol, which did not sufficiently
increase the viscosity of the solution inside the
thylcellulose, Int. J. Pharm. 67 (1991) 21–27. capsule to suspend the insoluble form of nifedipine $[4]$ G. Santus, R.W. Baker, Osmotic drug delivery: a review of so it could be released. Therefore, drug release the patent literature, J. Controlled Release 35 (1995) 1–21. operated by both osmotic and suspension mecha-

is not the development of osmotically controlled oral drug delivery

in the development of osmotically controlled oral drug delivery nisms. It should be pointed out that the sustainability
of drug suspension caused by the thickening effect of
 $\frac{1}{6}$ S.M. Herbig, J.R. Cardinal, R.W. Korsmeyer, K.L. Smith, the polymer was equally important as that of the Asymmetric membrane tablet coatings for osmotic drug delivery, J. Controlled Release 35 (1995) 127–136.

tures under a reasonable assumption that the active

period of the osmotic effect for both cases did not

significantly differ. The effect of solubility on the incorporation of solubility enhancers is able

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